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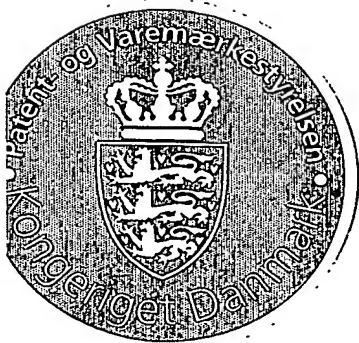
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Modtaget

Diagnostics of diarrheagenic *Escherichia coli* (DEC) and *Shigella* spp.

Field of invention

The present invention relates to a novel diagnostic assay for the detection of diarrheagenic *E. coli* (DEC) by identification of specific genetic markers, e.g. by use of multiplex PCR. The method further allows the evaluation of the pathogenic potential, which is valuable in relation to the treatment of a patient. The method will be useful for the analysis of any material from where alive bacteria can be generated, or from where bacterial DNA can be extracted. The specific PCR product can be detected by a number of technologies that are faster and both more sensitive and specific than conventional electrophoresis. The invention also includes a method for the subtyping a number the *E. coli* virulence genes that are believed to be important in the treatment and epidemiological surveillance of diarrheagenic *E. coli* infections.

General background

Diarrheagenic *E. coli* (DEC) strains isolated from intestinal diseases have been grouped into at least six different categories based on epidemiological evidence, phenotypic traits, clinical features of the disease they produce, and specific virulence factors. The currently recognized categories of diarrheagenic *E. coli* include: Attaching and effacing *E. coli* (A/EEC) including Enteropathogenic *E. coli* (EPEC), enterotoxigenic *E. coli* (ETEC), Enteroinvasive *E. coli* (EIEC), Enteroaggregative *E. coli* (EAaggEC), diffusely adherent *E. coli* (DAEC), and Shiga toxin-producing *E. coli* (STEC), which are also referred to as Verocytotoxin-producing *E. coli* (VTEC).

Table 1. Showing groups of diarrheagenic *E. coli* and characteristic virulence genes.

<i>E. coli</i> group	Positive for gene(s)	Corresponding toxin	Comments
VTEC	<i>vtx1</i> and/or <i>vtx2</i>	VT1 and/or VT2	May contain <i>eae</i> and/or <i>ehxA</i>
A/EEC	<i>eae</i>	Eae	Negative for any toxin genes and not belonging to the classical EPEC O:H serotypes.
EPEC	<i>eae</i>	Eae	Belong to classical O:H serotypes. Typical EPEC O:H serotypes are <i>bfpA</i> -positive, atypical EPEC are <i>bfpA</i> -negative. VTEC related EPEC strains may contain <i>ehxA</i> .
ETEC	<i>sta</i> and/or <i>elt</i>	ST and/or LT	
EIEC	<i>ipaH</i>	IpaH	

The most important groups are EPEC, ETEC, EIEC and VTEC whereas the role of EAggEC and DAEC are still being questioned. The definitions of these groups are not definitive and related to a number of genotypic- and phenotypic methods of characterization.

- 5 A definition adopted in 1995 identified the most important characteristics of EPEC as its ability to cause attaching and effacing (A/E) histopathology and its inability to produce Verocytotoxins. Typical EPEC of human origin possess a virulence plasmid known as the EAF (EPEC adherence factor) plasmid that encodes localized adherence on cultured epithelial cells mediated by the Bundle Forming Pilus (BFP), while atypical EPEC do not
- 10 posses this plasmid. The majority of typical EPEC strains fall into certain well-recognized O:H serotypes (10). According to this definition, the basic difference between typical and atypical EPEC is the presence of the EAF plasmid encoding BFP in the first group of organisms and its absence in the second. The definition is not static and may be changed as new types are discovered and described. The EPEC O:H serotypes that are currently regarded
- 15 as classical and newly recognised EPEC O:H serotypes by The International *Escherichia* and *Klebsiella* Centre (WHO) are shown in table 2.

Table 2. O:H serotypes regarded as classical and newly recognised EPEC O:H serotypes

O group	H antigen ^{a)}	Comments
O26	H ⁻ ; H11	O26:H ⁻ and O26:H11 may also be STEC/VTEC
O55	H ⁻ ; H6; H7	O55:H7, H10 and H ⁻ may also be STEC/VTEC
O86	H ⁻ ; H 8; H34	O86:H ⁻ may also be EAggEC H8 is a new type
O88	H ⁻ ; H25	New type
O103	H2	
O111	H ⁻ ; H2; H7	O111:H ⁻ may also be STEC/VTEC or EaggEC
O114	H ⁻ ; H2	
O119	H ⁻ ; H2; H6	
O125ac	H ⁻ ; H6	O125 may also be EaggEC
O126	H ⁻ ; H2; H21; H27	
O127	H ⁻ ; H6; H9; H21; H40	
O128ab	H ⁻ ; H2; H7; H12	O128:H2 may also be STEC/VTEC
O142	H ⁻ ; H6; H34	
O145	H ⁻ ; H45	New type
O157	H ⁻ ; H8; H16; H45	New types
O158	H ⁻ ; H23	

^{a)} Non motile strains of *E. coli* are regarded as descendants of motile strains that have lost their motility by mutation(s).

Apart from the well-recognized classical O:H EPEC serotypes, a large group of non-classical A/EEC serotypes of *E. coli* strains are found to be positive for the *eae*-gene. Together with EPEC, this group is referred to as Attaching and Effacing *E. coli* (A/EEC) based on the presence of the *eae*-gene and absence of toxin- or invasion genes. Like EPEC, they may be positive or negative for the EAF plasmid but they may also be positive for the *ehxA* plasmid found in many VTEC strains, see below.

ETEC strains do not invade epithelial cells but produce one or more enterotoxins that are either heat-labile (LT), which is closely related to cholera toxin, or heat-stable (ST).

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EIEC are very similar to *Shigella*. Like *Shigella*, they are capable of invading and multiplying in the intestinal epithelial cells of the distal large bowel in humans. Genes involved in the invasive phenotype of EIEC and most *Shigella* spp. are carried on a 140 MDa plasmid designated pInv. Prominent among these virulence genes is a type III secretion system (18). Also characteristic for the invasive phenotype is the *ipaH* gene, which is present in several copies on both the chromosome and the plasmid, making it especially suited as a diagnostic marker for EIEC and *Shigella* spp. (27).

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VTEC strains are characterized by their ability to produce either one or both of at least two antigenetically distinct, usually bacteriophage-mediated cytotoxins referred to as Stx1 or VT1 (first described as Shiga-like toxin I, SLTI) and Stx2 or VT2 (first described as Shiga-like toxin II, SLTII). Whereas STEC/VTEC refers to all *E. coli* strains that produce Stx/VT in culture supernatants (14,15), the term enterohemorrhagic *E. coli* (EHEC) has been used to refer to strains that have the same clinical and pathogenic features associated with the prototype organism *E. coli* O157:H7 (16). In practice, EHEC is used to describe a subgroup of STEC/VTEC that causes hemorrhagic colitis (HC). Almost all STEC/VTEC O157:H7 strains harbour a large 60-65 MDa plasmid (9), designated pO157, which plays a role in the virulence(11). The large plasmid of O157 encodes the EHEC-hemolysin (Ehx), which is homologous to the *E. coli* α -hemolysin (20,21). A role for Ehx in the pathogenesis of diarrhoeal disease has not been demonstrated but *ehxA* positive VTEC strains have been found more often in patients with Hemolytic Uremic Syndrome (HUS) than in patients with diarrhoea (6) and, together with the *eae*-gene in VTEC strains, serve as a predictor for more

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serious complications. O26:H11 strains also possess at least one plasmid in the range of 55-70 MDa and other O:H serotypes show a notable similarity with the large plasmids in O157 and O26 strains (16).

- 5 The diagnosis of DEC began in 1945 when Bray demonstrated the relation between *Bacterium coli var. neapolitanum* and diarrhoea in humans (3). A few years later Bray and Beaven used slide agglutination to type 95% of the bacteria isolated from stool cultures from children with diarrhoea (4). The breakthrough in typing was achieved in 1950, when the *E. coli* serotyping scheme was developed by Kauffmann (12). During the 1950s several new serogroups were
- 10 added to the list of those epidemiologically incriminated as causing diarrhoea (30). Meanwhile the enterotoxins of enterotoxigenic *E. coli* (ETEC) and the invasive properties of enteroinvasive *E. coli* (EIEC) had been described. Methods for detection included an infant mouse assay for the detection of ST, cell assays for LT, and inoculation of the eye of Guinea Pigs and subsequent development of keratoconjunctivitis for the detection of EIEC. In 1977, Konowalchuk *et al.* (15)
- 15 discovered a cytopathic effect in Vero cells from culture filtrates of *E. coli*. The effect could only be seen in Vero cells and not in Y1 mouse adrenal cells and Chinese hamster ovary (CHO) cells, and it was distinctly different from that of heat-labile enterotoxin. The cytotoxic effect was caused by one or more cytotoxins referred to as Vero toxins (VT) or Verocytotoxins (13).
- 20 Because of the above mentioned type diversity, rapid and easy methods of moderate cost for reliable identification and isolation of DEC strains independent of their serotype are required. A number of suitable methods for this purpose have been developed for each of the types but there is no internationally recognised standard procedure. These methods include biological assays, immunological methods, nucleic acid based assays or phenotypical tests such as O
- 25 grouping of commonly occurring DEC serotypes, enterohaemolysin production of the majority of VTEC types or the failure to ferment sorbitol or produce β -glucuronidase by most VTEC O157, and culture methods.
- 30 Unfortunately, these screening methods are incomplete because they are only directed against a subset of the DEC strains.

The diagnosis of DEC have important implications for the evaluation of possible intervention during the course of illness. Prolonged diarrhoea caused by EPEC and A/EEC especially in children may require antibiotic treatment of the patient whereas treatment of patients with a VTEC infection is not recommended due to the possible increased risk of a more severe outcome. In many countries, patients with a VTEC infection are quarantined or otherwise isolated due to the risk of contaminating other people. ETEC diarrhoea is not a very serious disease and usually self-limiting. It therefore usually does not require treatment. As is the case for VTEC infections, EIEC and *Shigella* infections are often succeeded by both quarantine and antibiotic treatment due to the low infectious dose and the risk of contamination other people.

Pass, M. A. et al. (2001) (19) have published a method for detection of pathogenic *E. coli* in cultured faeces. PCR was used to amplify specific fragments in the genes encoding the following 11 virulence factors: VT1, VT2, VT2e, CNF1, CNF2, LTI, STI, STII, EaeA, Einv and Eagg. It is stated that 4 multiplex-PCR combinations of primers gave adequate amplification of their respective genes. However, when the combination of multiplex-PCR with VT1, VT2, VT2e, EaeA, CNF2, Einv, LT and ST is shown, VT2e and VT1 are not visualised on the gel. This is an accepted fact that is explained in the article. Furthermore, the assay does not include a positive control.

Patent Application WO9848046 describes a PCR assay that detects EHEC, ETEC, EPEC, EIEC and EAggEC that are specifically designed for real-time PCR analyses. However, real-time PCR is presently limited to 4 simultaneous genes per reaction because of the fluorophore overlap.

López-Saucedo, C et al (2003) (17) are describing a method where the following 7 genes are detected in the same multiplex-PCR: *elt*, *sta*, *bfpA*, *eaeA*, *vtx1*, *vtx2* and *ial*. They are analysing the PCR products by size identification on agarose gel electrophoresis; the assay does not include the *ehxA* gene and does not have a positive control.

Compared to prior art the present method contains the following advantages:

- 5 - the use of *ipaH* for the detection of EIEC and *Shigella* spp. is a good genetic marker for this group of bacteria, as the gene is present in several copies both on the pInv plasmid and on the chromosome . The use of *ial* is a poor diagnostic marker for these bacteria because it is only present on the plasmid, which is easily lost both *in vivo* and *in vitro*.
- 10 - The use of *ehxA* as a diagnostic marker allows a further estimation of the pathogenic potential giving rise to serious diseases, which is not possible by any of the prior art.
- 15 - The use of 16S rDNA as positive control and the UNG system makes this method suitable as a reliable method for routine diagnostics. None of the prior art contains such considerations.
- 20 - The present method contains thoroughly validated tests both with respect to sensitivity and specificity (see example 1).

25 EPEC plasmids encoding the *bfpA*-gene and EHEC plasmids encoding the *ehxA*-gene have not been found together in the same strain and the two genes may therefore serve as useful genotypic markers for the presumptive categorization of any *eae*-positive *E. coli* as either belonging to the A/EEC - EPEC group or to the EHEC group.

30 The method disclosed in the present invention detects the clinically relevant DEC types simultaneously and has very few limitations. The big advantage of this invention is that subsequent to the identification of positive stool cultures, procedures for further analysis by supplementary PCR of the bacterial lysates obtained during screening are possible and could include: virulence gene subtyping by PCR followed by restriction digests or sequencing, O:H serotyping by sequencing of PCR amplified bacterial antigens, or other genotyping by for example microarray analyses. The procedure also allows for the referral of the lysate to more specialised reference laboratories, which – in times where bioterrorism is ever threatening - will be safe and easy to understand for everybody at the primary screening laboratory facility. All primers chosen in the present invention were designed to match the most conserved

regions within the relevant genes. By doing so, the method is optimised to detect any possible subtype of the relevant genes, including new genetic subtypes that are expected to contain genetic changes in the less conserved regions, increasing the chance of being detected by the present method. However, as for any PCR based method, it requires continuously updating and validation whenever new genotypes are being described. As our laboratory serves as The International *Escherichia* and *Klebsiella* Centre (WHO) there will be no problem in obtaining presumptive new types.

10 Summary of the invention

The presently preferred embodiments of the present invention are outlined in the following points:

- 1) Novel multiplex-PCR combination detecting the 8 genes *bfpA*, *ehxA*, *vtx1*, *vtx2*, *eaeA*, *ipaH*, *sta* and *elt*, which is found to be the most suited gene combination for the characterization of diarrheagenic *E. coli*., and including a PCR-control derived from 16S rDNA (positive control),
- 2) Intensive validated experimental procedure, showing superior sensitivity and specificity compared to other publications.
- 3) Descriptions of how the multiplex-PCR can be combined with other technologies in order to decrease time of analysis and improve sensitivity.
- 4) Routine diagnostic consideration with respect to carry-over prevention, by the use of the UNG system.
- 5) Protocols for the subtyping of the virulence genes: *eae*, *vt1* and *vt2* by either, direct sequencing of the amplicons generated in the multiplex-PCR, or by sequencing of a larger fragment generated by a new PCR.

Detailed description of the invention

The present invention discloses a method for simultaneous detection of diarrheagenic *Shigella spp.* and *E. coli* (DEC) including A/EEC & EPEC, ETEC, VTEC, EIEC and especially strains with the *ehxA* gene.

A preferred embodiment of the invention detects the presence of the genes *ehxA*, *eae*, *vtx1*, *vtx2*, *ipaH*, *stx*, *elt* and *bfpA*. and is able to detect *Shigella spp.* by the presence of the *ipaH* gene. The presence of the genes can be detection of the genes themselves or parts herof, RNA or polypeptides coded by the genes.

The screening method can be performed by nucleotide sequence amplification technique, such as PCR, multiplex PCR, real-time PCR, most preferably multiplex PCR, with a selected set of primers and incorporating a positive control using 16S rDNA. Possible contamination of samples are preferably reduced by incorporating the UNG system.

Detection of the genes can be performed by size identification, e.g. by agarose gel electrophoresis or capillary electrophoresis or with a hybridisation probe.

The sample material to be analysed can be any material from where bacteria can be extracted, e.g stool samples, consumables etc.

The screening method can be used as an *in vitro* diagnostic method for determining the risk of being infected with a pathogenic organism which gives rise to haemolytic uremic syndrome (HUS) or hemorrhagic colitis, by detecting the *ehxA* gene in the sample.

The invention discloses a specific set of primers and probes for the respective genes but are not restricted to these.

The invention also discloses a kit for the screening, which comprises, in a single or in separate containers, nucleotide sequences which are able to prime amplification in a nucleotide sequence amplification reaction, such as PCR, of the genes: *ipaH*, *eae*, *ehxA*, and

st, or parts of these genes or the complementary strands to the genes or parts thereof.

Additionally the kit can comprise nucleotide sequences which are able to prime amplification in of the genes: *vtx1*, *vtx2*, *elt*, and *bfpA*, or parts of these genes or the complementary strands to the genes or parts thereof. The described kit can also contain nucleotide sequences which
5 are able to hybridise (preferably under stringent conditions) with the genes: *ipaH*, *eae*, *ehxA*, *sta*, *vtx1*, *vtx2*, *elt*, and *bfpA* parts of these genes or the complementary strands to the genes or parts thereof. Preferably the kit comprises a means for a control, such as primers for 16S rDNA.

10 Definitions and abbreviations:

A/E: attaching and effacing

A/EEC: attaching and effacing *E. coli*

Amplicon: *syn.* "PCR product"

bfpA: bundle forming pilus, structural gene, subunit A; Virulence factor in EPEC, which is involved in the initial adherence of the bacteria to the intestinal cells.

bp: base pair

Capillary electrophoresis: Capillary electrophoresis (CE), is a technique where an electrophoretic separation takes place in a thin capillary tube filled with buffer. A sample is injected at one end, either by electrophoresis or by pressure, and an electric field of 100 to 700 volts/centimeter is applied across the capillary. It is generally used for separating ions, which move at different speeds depending on their size and charge, when the voltage is applied. At the end of the capillary each of the separated analytes are measured by a detector in a time dependent manner. CE is usually run with an internal standard that allows size determination of the separated sample molecules.

Carry-over prevention: dUTP is incorporated in all PCR products instead of dTTP. Before PCRs are subjected to thermocycling, they are incubated with UNG (uracil-DNA glycosylase) that

degrades any single or double stranded DNA containing dUTP, but has no effect on dTTP containing DNA. By this procedure, possible contamination from other PCRs is reduced, while the amplification of bacterial DNA is unaffected.

DAEC: diffusely adherent E. coli

DEC: diarrheagenic E. coli

ea: E. coli attaching and effacing: intimin. Virulence factor from EPEC or VTEC.

EAF: EPEC Adherence Factor plasmid. Plasmid containing BFP

EaggEC: enteroaggregative E. coli , syn. EAEC

EHEC: enterohemorrhagic E. coli

ehxA: enterohemolysin, structural gene, subunit A

EIEC: enteroinvasive E. coli

elt: gene encoding heat labile enterotoxin (LT)

EPEC: enteropathogenic E. coli

ETEC: enterotoxigenic E. coli

GI: gastrointestinal tract

HC: hemorrhagic colitis

HUS: haemolytic uremic syndrom

ipaH: invasive plasmid antigen H

LT: heat-labile enterotoxin

Luminex technology: microbeads of different internal colors are labeled with hybridization probes representing different genes. These beads are hybridized with fluorescence labeled sample DNA (usually PCR products) under stringent condition. After the hybridization has taken place, the mixture is injected into the instrument that uses microfluidics to align the microbeads in a single file where lasers illuminate the colors inside and on the surface of each microbead. The optics capture the combination of color coded microbeads and hybridized sample molecule.

Microarray technology: hybridisation probes representing different genes are chemically linked to different spots on a solid surface, usually a small glass slide. Fluorescence labeled sample DNA (usually PCR products) are hybridised to the capture probes under stringent condition. After the post hybridization washing steps, only sample DNA with nucleotide sequences complementary to the capture probes will stay bound to the slide. Bound PCR products at specific spots of known capture probes, are registered by their fluorophore emission.

Multiplex PCR: PCR with more than one primer set present in the same reaction, where each primer set is amplifying a unique locus if the specific template is present.

O:H: specific serotype; "O" refers to the LPS O antigen, and H refers to flagellar antigen.

Real-time PCR: Detection of the PCR while the temperature cycling is still in progress. This can be done by the measurement of emitted fluorescence, which is linked to the reaction. The fluorescence can originate from fluorophores that bind to the double stranded amplicons (sequence unspecific intercalating agent, ex; SYBR Green), or it can originate from sequence specific probes that are designed downstream of the primers. Such probes will emit light only when digested by the polymerase because they contain a fluorophore and a corresponding quencher. Real-time PCR is limited to 4 simultaneous genes per reaction because of the fluorophore overlap.

stx 1 / 2: genes encoding verocytotoxin 1 / 2

ST: heat stable enterotoxin

sta: gene encoding heat stable enterotoxin (ST)

STh: heat stable enterotoxin (human), *syn.* STIB

STp: heat stable enterotoxin (porcine), *syn.* STIA

VT 1 / 2: verocytotoxin 1 / 2, *syn.* shiga like toxin (Stx)

VTEC: verocytotoxin producing *E. coli*

vtx 1 / 2: gene encoding verocytotoxin 1 / 2 (VT 1 / 2)

UNG: uracil-DNA glycosylase

Enclosed in the present invention is the possibility of performing diagnostic PCR, both on DNA prepared directly from human faecal samples, or on DNA prepared from colonies
5 grown from faecal samples. Performing PCR on DNA purified directly from faeces is an attractive strategy, as it saves time and labour. Besides that, direct PCR can detect dead cells, and cells prone to loose plasmids during *in vitro* growth, and is not affected by the selectivity that a growth step might introduce. However, it is also important to have a fast and easy way to test plated out cell material, when single colonies need to be isolated. For that reason, the
10 present invention contains a method, which relies on simple boiling and centrifugation for preparing PCR-usable template DNA.

In the present invention, sensitivity limits are established by making a dilution series of pathogenic bacteria in a background of non-pathogenic bacteria. That is the most important
15 way to test the sensitivity, as this situation is closest to the compositions of clinical samples. In the present invention, a positive result can be obtained, if the template DNA has a composition, where one pathogenic bacterium is present among 10^4 non-pathogenic bacteria.

High specificity relies on good primer design, but is also depended on experimentally testing of the assay on different strains know to harbour different homologous of the relevant genes. The present invention has been tested 100% specific on 116 different reference strains
5 obtained from The International *Escherichia* and *Klebsiella* Centre, WHO, Statens Serum Institut, Denmark (see table 5).

10 In a presently preferred embodiment, the present invention is using the genes coding for the following virulence factors: BfpA, EhxA, VT1, VT2, Eae, IpaH (same as Einv), ST and LT. Genes encoding both VT2, VT2c, VT2d and subtype VT2e will be amplified by the present PCR, and result in a PCR product of the same size. Besides that, the present invention has included a universal primer-pair towards 16S rDNA as a positive control for the PCR. As for
15 the combination of genes in the multiplex-reaction, the present invention is able to perform sensitive and specific PCR with all 8 virulence genes and 16S rDNA present in the same multiplex reaction.

The present invention includes all the relevant genes for the currently recognised and clinically important groups of DEC. The choice of genes allows for a rapid evaluation of the
20 further treatment and interventional strategies in relation to the individual patient in order to minimise complications and the spread of highly pathogenic bacteria to contacts or the environment.

The present invention contains a method for the subtyping of the virulence factors VT1, VT2, Eae and other genes. Subtyping of these virulence genes is increasingly being accepted as an
25 important part of characterizing VTEC infections, especially in relation to the proper treatment (5).

The present invention solves the diagnostic problem of screening for human pathogenic *E. coli* groups. The method relies on specific multiplex-PCR amplification of 8 virulence genes
30 allowing a distinction between the pathogenic *E. coli* groups: ETEC, VTEC, A/EEC including EPEC and EIEC, and provides important distinction between typical/atypical EPEC strains and additional information on the presence or absence of the EHEC plasmid.

The method is based on primers chosen to match all clinically relevant subtypes of the given virulence genes. The PCR setup is designed to enclose all primer sets in one single reaction, leading to the specific amplification of any given template present. The method was optimised to result in the best sensitivity and specificity. This was done by analysing DNA from 10-fold serial dilutions of bacterial colonies known to harbour different subtypes of the relevant virulence genes. The method therefore, allows for the analysis of any material from where alive colonies can be cultured. Of special interest is, the analysis of stool samples from diseased patients, where time, sensitivity and specificity are critical parameters in effective diagnoses. Also, the analysis of consumables has a high value as it may prohibit the spread and intake of contaminated foods. Due to the well characterized sensitivity limits, the methods also has the potential of analysing DNA purified directly from the primary sources.

The PCR primers, -probes, -reagents and -temperature conditions were optimised to perform well in combination with a number of different technologies. These technologies fall into two different groups

- DNA purification directly from the source. This can be done by a number of different commercial kits described in section 5 (Example 3-6).
- PCR products detection by either capillary electrophoresis, real time PCR or solid-face capture probe techniques like; membrane/ELISA hybridisation, DNA chips or Luminex.

The present invention also encloses a real-time PCR setup with optimised PCR conditions including specific primer and probe design. Besides that, the present invention also contains the option of amplifying all 9 genes in the same reaction, in a both sensitive and specific manner (non-real-time PCR setup). This is possible because the concentration of every reagent has been carefully optimised, and because the present invention (in this setup) is not burdened by the addition of a specific probe for every gene in the assay.

The present invention includes hybridisation probes specifically designed and optimised for constituting the capture probes, in solid surface hybridisations like membrane hybridisation or hybridisation in microtiter plates, DNA microarrays and hybridisation on microbeads (ex. Luminex ® technology). Finally, the PCR products of the present invention lie within the

size-range that should be easily detectable by capillary electrophoresis, which is faster, more sensitive and accurate than gel-electrophoresis.

Being able to use these technologies in combination with the multiplex PCR, results in a number of advantages compared to traditional diagnostics. Firstly, direct DNA purification from the source is not affected by the selectivity that a growth step might introduce, dead bacteria and bacteria that easily loose plasmids can be detected, and the entire procedure is much faster. Secondly, due to the multiplex setup, it only requires one PCR to screen for the entire 8 virulence genes. Thirdly, the technologies used for amplicon detection are faster and more sensitive than traditional methods.

Specific primers are of major importance in a diagnostic PCR setup. The pivot of this problem is sequence analysis of the available data in Gene Bank. The primer and probes were designed on basis of the considerations described below.

The heat stable enterotoxin (ST) of ETEC is a small monomeric protein of 18-19 amino acids encoded by the *sta* gene which is 219 bp long (18). The relative few submitted nucleotide sequences of *sta*, available in Gene Bank, fall into a number of phylogenetic groups based on ClustalW comparisons. The groups of genes encoding heat stable enterotoxins were as follows: five *staI* genes (accession numbers: J03311, M34916, M29255, M18346 and M18345) made up their own cluster, two other *staI* genes (accession numbers: M25607 and M58746) were more related to the heat stable enterotoxins from *Yersinia enterocolitica* and *Vibrio cholera*, and two separate clusters were made up of *staII* genes and the gene encoding EAST1. As the 5-gene cluster of *staI*, seems to be the major heat stable enterotoxin produced by human ETEC strains (18) (22), and as their sequences were too divergent from the other heat stable enterotoxins, probe and primers were designed towards those sequences only. When the primer-set was tested on *Yersinia enterocolitica*, *Vibrio cholera* and different *E. coli* strains (listed in table 5) 100% specificity was observed for the aforementioned *staI*. The primers were chosen to amplify a 170 bp fragment.

The heat labile enterotoxin (LT) of ETEC is composed of one A-subunit and 5 identical B-subunits encoded by the *elt* gene. The toxin can be divided into LTI and LTII based on

serology and host pathogenesis. The genes encoding the A- and B-subunits are 777bp and 375bp long, respectively (7). The cholera enterotoxin produced by *Vibrio cholerae* is about 75% identical in the nucleotide sequence to the LT1 of *E. coli*. The sequences of subunit A from nine *eltI* genes (accession number: V00275, S60731, AF242417, AB011677, M35581, M15261, K01995, M15362 and M57244) were compared to a number of *eltIII* and *Vibrio cholerae* *ctx* genes. Due to the desired specificity towards *E. coli*, the clinical unimportance of *eltIII* (18) and the relative low homology between *eltI* and *eltIII* / *ctx*, probe and primers were designed to match *eltI* only, and result in an amplicon of 479 bp.

Intimin is encoded by the *eae* gene in either A/EEC (including EPEC) or VTEC, and has a size of approximately 2810 bp. Based on the divergent sequences in the last third of the 3-prime end, at least 8 subtypes can be identified. At least one of each subtype was present in the gene alignment and the following accession numbers were used: AF081186, AF253560, U60002, AB040740, AJ308552, AF116899, AF449419, AF081184, AJ308551, AF449416 and AJ298279. Probe and primes were designed to match all tested gene sequences, and the PCR-product has a size of 377 bp.

The virulence factor, bundle forming pilus (Bfp) from EPEC, is encoded by an operon consisting of 14 genes, including the 580 bp structural gene *bfpA*. Based on sequence comparisons, the *bfpA* genes fall into an alpha and a beta type. Probe and primers were designed to match both types by aligning the genes with the following accession number: AF304478, AF304486, AF304482, AB024946, AF304480, AF304477, AF304484, Z12295, AF382948. The resulting amplicon was 307 bp long.

Both vero toxin 1 and 2 (VT1 and VT2) from VTEC are composed of an A- and a 5 B-subunits. The gene encoding the A-subunit is approximately 960 bp long, and the gene encoding the B-subunit is approximately 270 bp long. As the homology between VT1 and VT2 is relatively low (about 50% identity), and because of the desirable differentiation between the two toxins, specific probes and primers were designed to each gene.

Based on the nucleotide sequence it is difficult to distinguish between *vtx1* from *E. coli* and shiga toxin 1 from *Shigella* spp. Also, all *vtx1*-genes from *E. coli* share very high homology. Probe and primers were designed to match all *vtx1*- genes from *E. coli* and all shiga toxin 1 genes from *Shigella* spp, by alignment of the genes with the accession numbers: AF461172, AJ279086, AF153317, AJ132761, Z36899, AB030485, AB035142, AF461166, AJ251325, AJ314839 and M19473. The resulting PCR product is 260 bp long.

Most *vtx2*-genes share relative high identity (above 90%). However, one group of genes seems to make a unique cluster, consisting of the *vtx2f* (accession numbers AJ270998 and AJ010730) and *vtx2va* (M29153) (now renamed *vtx2f*) subtypes, with about 60% identity to the other *vtx2* genes. Due the relative low sequence homology, and the fact that most *vtx2f* and *vtx2va* are not found in humans (5), probe and primers were designed to match the major *vtx2* group, by aligning the *vtx2* sequences: AJ313015, AP000363, AB048228, L11078, X81415, X81418, X61283, M36727, AB017524, AF291819 and Y10775. The PCR product for VT2 was designed to be 420 bp long.

Enterohemolysin A (Ehx), often found in EHEC is encoded by the ca. 3000 bp long *ehxA* gene, which is part of the 4-gene enterohemolysin operon. The *ehxA* genes are a very homogenous group of genes, and probe and primers were designed to match all known subtypes. The following accession numbers were used in the gene alignment: X79839, AB032930, AF074613, X86087, X94129, AB011549 and AF043471. The PCR product for *ehxA* was designed to be 530 bp long.

The invasive plasmid antigen *ipaH*-gene is carried in multiple copies on both the 140 MDa invasive plasmid as well as on the chromosome of EIEC strains and *Shigella* spp. The advantage of using this gene, rather than the *ial*-gene is that it remains detectable despite the loss of the plasmid. Probe and primers were designed to match all genes analysed in the gene alignment, made up of the following genes: AL391753, M32063, AF047365, M76445, M76443, AF386526, AF348706 and M76444. The size of the PCR product is 647 bp.

16S rDNA was chosen as a positive control, because many Gram-negative bacteria found in the human GI share high sequence homology. Thus, the detection of templates, that matches

the primers and probe is very high under any given clinical conditions (even in antibiotic treated patients). Probe and primers were chosen to match as many as possible of the common bacteria from the human GI. The 16S rDNA control band is 1062 bp long.

5 Table 3. Listed are the 8 virulence genes and 16S rDNA used in the multiplex-PCR assay, together with amplicon size, final primer concentration, and primer sequences.

Virulence factor	Gene locus	E. coli group	Primer-name	Amplicon size (bp)	Primer conc. (μM)	Primer sequence (5'→3')
Heat labile enterotoxin I (LT1)	<i>elt</i>	ETEC	LT-F	479	0.4	AAACCGGCTTTG-TCAGATATGATG
			LT-R		0.4	TGTGCTCAQAT-TCTGGGTCTCC
Heat stable enterotoxin I (ST1)	<i>sta</i>	ETEC	ST-F	171	0.25	TCACCTTTTCG-CTCAGGATGC
			ST-R		0.25	ATAGCACCCG-GTACAAGCAGG
Intimin (Eae)	<i>eae</i>	EPEC /	eae-F	377	0.2	GGYCAGCGTT-TTTTCCTTCCTG
		VTEC	eae-R		0.2	TCGTCACCAR-AGGAATCGGAG
Shiga toxin I (Stx1) / Verocytotoxin I (VT1)	<i>stx1</i> /	VTEC	VT1-F	260	0.25	GTTTGCAGTTG-ATGTCAGAGGGA
	<i>vtx1</i>	Shigella	VT1-R		0.25	CAACGAATGG-CGATTTATCTGC
Shiga toxin 2 (Stx 2) / Verocytotoxin 2 (VT2)	<i>stx2</i> /	VTEC	VT2-F	420	0.25	GGAATGCAAATC-AGTCGTCACCTC
	<i>vtx2</i>		VT2-R		0.25	GCCTGTCGCCA-GTTATCTGACA
Invasion plasmid antigen H (ipaH)	<i>ipaH</i>	EIEC /	ipaH-F	647	0.1	TTGACCGCCT-TTCCGATACC
		Shigella	ipaH-R		0.1	ATCCGCATCA-CCGCTCAGAC
Enterohemolysin A (EhxA)	<i>ehxA</i>	VTEC /	ehx-F	533	0.05	GGGAAAAGCC-GGAACAGTTCT
		EPEC	ehx-R		0.05	CCAGCATAAC-AGCCGATGTGAT
bundle-forming pilus A (BfpA)	<i>bfpA</i>	EPEC	bfp-F	307	0.4	TCCAATAAGKC-GCAGAATGCTA
			bfp-R		0.4	CACCGTAGCCT-TTCGCTGAAG
16S rDNA	16S	most	16S-F	1062	0.25	GGAGGCAGCA-GTGGGGAATA
		gram +	16S-R		0.25	TGACGGGCGG-TGTGTACAAG

R = G or A, Y = C or T, K = G or T

10 Primers were designed with the following parameters: 55-57°C melting temperature, GC-content between 45-60%, length between 20-24 bp, lowest possible likelihood of dimer- and hairpin formation, optimal entropy in the ends and distinguishable amplicon sizes. Each primer set was tested individually under varying PCR conditions, and the optimum conditions were used to construct the PCR conditions in the multiplex reaction. Primers were redesigned until they met the desired level of sensitivity and specificity, when all 9 primer

sets were able to amplify any given target present in the sample. Probes were designed to have a melting temperature between 65°C and 67°C, the least possibility of dimer- and hairpin formation and no G in the 5'-end. The primer sets were optimised by testing the PCR methods on a number of different strains under different template conditions.

5 The present invention also contains a method for subtyping the *vtx1*, *vtx2* and the *eae* genes. By using one of the PCR primers in a sequencing reaction, performed on the PCR products, the resulting sequence can be phylogenetically analysed and assigned to a specific subtype by the comparison to sequences of known subtype. If the PCR product does not contain subtype specific sequence, a set of PCR primers can be designed to amplify a larger fragment
10 containing more subtype specific sequences.

The present invention contains, PCR conditions that have been optimised to work with the carry-over prevention systems using dUTP and UNG. In order to prevent contamination from other PCRs, dUTP is incorporated into PCR products instead of dTTP. Before PCRs are
15 subjected to thermocycling, they are treated with UNG that degrades dU-containing PCR products, but has no effect on native template DNA. The same level of sensitivity and specificity as described above is obtained when UNG and dUTP are included in the assay.

The PCR products can be analysed by size identification on agarose gel electrophoresis as every PCR-product has a unique size. Besides that, the present invention encloses a number
20 of faster and more sensitive solutions for identification of the PCR products. For each gene, a hybridisation probe has been designed, from a conserved region within the gene. This feature adds an extra level of specificity, as the PCR product must have the right internal sequence in order to be detected. The specific hybridisation probes can be utilized in a number of
25 different technologies. Firstly, real-time PCR can detect a positive PCR, before the thermocycling has run to completion. Besides the obvious time saving advantage, this technology is far more sensitive and specific than agarose gel electrophoresis. Real-time PCR is technically limited to a maximum of 4 genes per multiplex reaction, which means that the total of 9 genes needs to be analysed in 3 reactions. Secondly, the specific probes can
30 constitute the capture-probes on solid surfaces like nylon membranes, ELISA-plates or DNA microarrays, where a stringent hybridisation can take place, which is subsequently analysed due to colour-coded reagents. Thirdly, capture-probes can also be situated on microbeads (ex.

Luminex ® technology), where the specific hybridization is analysed from the combination of colour-coded beads and colour-coded PCR-products. Finally, and not based on hybridisation, PCR-products can be identified by capillary electrophoresis, which is faster, more sensitive and accurate than gel-electrophoresis.

5

Figure legends

Fig. 1. Two *E. coli* pathogenic reference strains are mixed in equal volumes, and serially diluted in a constant background of a non-pathogenic *E. coli* strain (D2103). One pathogenic *E. coli* strain harbours *vtx1*, *eae*, *vtx2* and *ehxA* (strain D2164), while the other pathogenic strain (fr1368) contains *ipaH*. All PCRs contain a total amount of DNA corresponding to the preparation of approximately 0.05 bacterial colony.

Lane 1: only D2103,

15 Lane 2 only D2164 and fr1368,

Lane 3: 1/10 of D2164 and fr1368 relative to D2103,

Lane 4: 1/102 of D2164 and fr1368 relative to D2103,

Lane 5: 1/103 of D2164 and fr1368 relative to D2103,

Lane 6: 1/104 of D2164 and fr1368 relative to D2103,

20 Lane 7: 1/105 of D2164 and fr1368 relative to D2103 and

Lane 8: 1/106 of D2164 and fr1368 relative to D2103

Fig. 2. Two *E. coli* pathogenic reference strains are mixed in equal volumes, and serially diluted in a constant background of a non-pathogenic *E. coli* strain (D2103). One pathogenic *E. coli* strain harbours *bfpA* and *eae* (strain D1826), while the other pathogenic strain contains *sta* and *elt* (strain D2168). All PCRs contain a total amount of DNA corresponding to the preparation of approximately 0.05 bacterial colony.

Lane 1: only D2103,

30 Lane 2 only D1826 and D2168,

Lane 3: 1/10 of D1826 and D2168 relative to D2103,

Lane 4: 1/102 of D1826 and D2168 relative to D2103,

Lane 5: 1/103 of D1826 and D2168 relative to D2103,
Lane 6: 1/104 of D1826 and D2168 relative to D2103,
Lane 7: 1/105 of D1826 and D2168 relative to D2103 and
Lane 8: 1/106 of D1826 and D2168 relative to D2103

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Fig. 3. PCR test of 8 different strains (lane 1-8). Four strains (lane 1, 2, 4, and 6) harbouring a combination of the 6 virulence genes: *sta*, *vtx1*, *eae*, *elt*, *ehxA* and *ipaH*. Lane 3, 5, 7 and 8 were tested negative for pathogenic *E. coli*. Lane 9 contains 100 bp DNA marker. Marker to the left of lane 1 was removed for better visualization of gene designation, but was present when genes were assigned.

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Fig. 4. PCR test of 8 different strains (lane 1-8) harbouring a combination of the 5 virulence genes: *vtx1*, *bfpA*, *eae*, *vtx2* and *ehxA*. Lane 9 contains 100 bp DNA marker. Marker to the left of lane 1 was removed for better visualization of gene designation, but was present when genes were assigned.

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Examples

Examples 1-2 contains experimental data that shows how this method perform with respect sensitivity and specificity, and how is can be applied as a diagnostic tool in a routine laboratory. Examples 3-6 are intended to illustrate the invention, and describe how is can be applied in combination with other technologies. The combination of the following steps: a) DNA extraction; b) multiplex PCR and; c) method of PCR product detection, is not the restricted to the ones mentioned in the examples, but will work in any preferred combination. The specific method was first developed to diagnose human *E. coli* infections from the bacterial presence in stool samples, but most of the DNA purification methods will work on a variety of different starting materials. Example 6 deals exclusively with the genetic subtyping of genes encoding either VT2 and/or *eae* from either VTEC or EPEC infections.

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Contents:

- Example 1:**
- Template DNA prepared from plate grown cells by simple boiling procedure.
 - Multiplex-PCR on the genes encoding the following *E. coli* virulence factors: ST,
5 LT, Eae, BfpA, VT1, VT2, EhxA and IpaH.
 - Identification of PCR products by gel electrophoresis.
 - This example shows how the multiplex-PCR method performs with respect to sensitivity and specificity.
- Example 2:**
- DNA extraction from bacterial colonies, derived from fecal samples by growing
fecal samples overnight on agar plates.
 - Multiplex PCR on the genes encoding the following *E. coli* virulence factors: ST,
15 LT, Eae, VT1, VT2, and IpaH.
 - Identification of PCR products by gel electrophoresis.
 - This example shows how a method performs in a routine diagnostic laboratory compared to a DNA hybridisation technique.
- Example 3:**
- DNA purified directly from feces by performing cell lysis and separation of magnetic beads that bind DNA (Kingfisher, Thermo Labsystems, Finland).
 - Multiplex PCR on the genes encoding the following *E. coli* virulence factors: ST,
20 LT, Eae, BfpA, VT1, VT2, EhxA and IpaH.
 - Detection of PCR products by LUMINEX™ technology.
 - This example describes a theoretical procedure for the above technologies.
- Example 4:**
- DNA purified directly from feces, by separation of magnetic beads that bind bacterial cells followed by cell lysis and ethanol wash (Genpoint A.S,
30 Norway)
 - Multiplex PCR on the genes encoding the following *E. coli* virulence factors: ST,

LT, Eae, BfpA, VT1, VT2, EhxA and IpaH.

- Detection of PCR product by hybridisation to solid-phase capture-probes on ex.

nylon membrane, plastic surfaces or DNA chip/microarrays.

- - This example describes a theoretical procedure for the above technologies.

Example 5: - DNA purified directly from feces, by DNA absorption/entrapment in fibrous membranes (FTA Technology, Promega)

- Multiplex PCR on the genes encoding the following *E. coli* virulence factors: ST,

LT, Eae, BfpA, VT1, VT2, EhxA and IpaH.

- Detection of PCR products by real-time PCR.

- This example describes a theoretical procedure for the above technologies.

Example 6: - DNA purified directly from feces by use of cell lysis, absorption and elution of

DNA from spin columns (ex. QIAamp® DNA Stool Mini Kit, QIAGEN).

- Multiplex PCR on the genes encoding the following *E. coli* virulence factors: ST,

LT, Eae, BfpA, VT1, VT2, EhxA and IpaH.

- Detection of PCR products by use of capillary electrophoresis

- This example describes a theoretical procedure for the above technologies.

Example 7: -Subtyping of the virulence genes *vtx2* and *eae*, by direct sequencing of amplicons

containing subtype distinguishable sequences.

Example 1:

INTRODUCTION

The present example describes the use of multiplex-PCR in the identification of the *E. coli* groups: ETEC, EPEC and A/EEC, VTEC and EIEC. The PCR relies on the specific multiplex-amplification of the genes encoding the following virulence factors: heat-labile enterotoxin (LT) characteristic for ETEC, heat-stable enterotoxin (ST) characteristic for ETEC, intimin (Eae) characteristic for EPEC or VTEC, bundle-forming pilus (BfpA) characteristic for EPEC, enterohemolysin (EhxA) characteristic for VTEC, vero toxin 1 and 2 (VT1 and VT2) characteristic for VTEC and invasive plasmid antigen (IpaH) characteristic for EIEC. As a positive control for the PCR a primerset for the 16S rDNA gene from most gram-positive bacteria was also included in the assay. The PCR analysis is performed on template DNA derived from bacterial colonies grown on plates. Each PCR was analyzed by agarose gel electrophoresis for the possible presence of amplicon(s) of the size that could be identified as any of the virulence markers mentioned above. The present example shows how the PCR method performs with respect to sensitivity and specificity. This includes the analysis of serially diluted reference strain and analysis of a 116 reference strains. All reference strains were also tested by a probe hybridization technique directed towards the same genes.

MATERIALS AND METHODS

Membrane hybridisation:

Each virulence gene that were screened for were contained on a specific pBluescript clone. These clones served as templates for the labeling reaction using the PCR DIG Labeling Mix from Roche, and T3/T7 pBluescript primers or primers designed within the gene. Strains used to construct the clones were obtained from The International *Escherichia* and *Klebsiella* Centre, WHO, Statens Serum Institut, Denmark and genes encoding the following factors were included in the assay: Eae (8), VT1 (28), VT2 (26), STp/ST1A (24), STh/ST1B (24), LT (24) and ipaH (27).

By use of a 1 µL sterile loop, small volumes of the different colonies, that were chosen to be investigated, were transferred to individual spots on a nylon membrane (Hybond-N⁺, Amersham Biotech) that was positioned on top of an agar plate. After over night growth the

nylon membrane was removed from the agar plate. The colonies on the nylon membrane were lysed, denatured and neutralized by incubating the nylon membrane in the following solutions for 10 min each: 10% SDS, 0.5 N NaOH, 1.4 M NaCl/0.5 M NaOH, 1 M Tris-HCl, pH 7.4 and 1.5 M NaCl / 0.5 M Tris-HCl, pH 7.4. The membrane was then baked at 65°C for one hour and prehybridised in 0.1 x SSC and 0.1% SDS at 65°C.

The probe was denatured by boiling in hybridisation solution for 8 min (0.5% Blocking Reagent, 0.1% N-laurylsarcosine, 0.02% SDS and 5 x SSC). The pre-hybridisation solution was exchanged with the hybridisation solution containing the denatured probe and incubated at 65°C for one hour. The hybridisation solution was discharged and the membrane was washed twice in 2 x SSC, 0.1% SDS for 5 min at room temp, and twice in 0.1 x SSC, 0.1 % SDS for 30 min at 65°C.

The hybridisation signals were developed by washing the membrane in Detection Buffer (0.1 M maleic acid, 0.15 M NaCl, 0.2 M NaOH, pH 7.5) for 2 min at room temperature, and in Blocking Buffer (1% Blocking Reagent, 0.1 M maleic acid, 0.15 M NaCl, 0.2 M NaOH, pH 7.5) at room temperature for 25 min. Next, 6µL Anti-digoxigenin was mixed with 60mL Blocking Buffer and added to the membrane and incubated at room temperature for 30 min. The membrane was then washed twice in Detection Buffer for 15 min each, and incubated for 2 min with Detection Buffer containing 50 mM MgCl₂.

Finally the membranes were incubated over night in the dark at room temperature in Detection Buffer containing 50 mM MgCl₂, 0.15% NBT (4-Nitro blue tetrazolium chloride, Boehringer Mannheim) and 0.1% BCIP (x-phosphate/5-Bromo-4-chloro-3-indolyl-phosphate, Bohringer Mannheim). Next day, the colour reaction is stopped, by rinsing the membrane in water, and the resulting hybridisation reactions were visualized for the individual spots.

PCR:

Relevant subtypes of each gene were downloaded from Gene Bank. Alignments (ClustalW algorithm) of gene sequences and primer design were done using the LaserGene Software DNASTar. Homologues of each gene that were included in the alignments are described above.

Primers sequences were chosen to have comparable GC content (45-60%), base pair length (20-24 base pairs) and melting temperatures (55-57°C), optimal 3' end and 5' end stability, and low likelihood of hairpin loop and primer-dimer formation.

Primer-sets based on the theoretical primer design, were tested experimentally at different annealing temperature and different concentrations of PCR reagents. The resulting optimum conditions were taken into account when the primer sets were combined in the multiplex analysis. Primers were redesigned until they met the satisfactory level of sensitivity and specificity. Each primer set was chosen to result in a unique amplicon size that could be easily identified by agarose gel electrophoresis. See table 3 for primer sequences and amplicon sizes.

Genes encoding the following virulence factors were included in the assay: heat-labile enterotoxin (LT) characteristic for ETEC, heat-stable enterotoxin (ST) characteristic for ETEC, intimin (Eae) characteristic for A/EEC, EPEC or VTEC, bundle-forming pilus (BfpA) characteristic for EPEC, enterohemolysin (EhxA) characteristic for VTEC, vero toxin 1 and vero toxin 2 (VT1 and VT2) characteristic for VTEC, and invasive plasmid antigen (IpaH) characteristic for EIEC. As a positive control for the PCR, a primer set targeting 16S rDNA matching most gram-negative bacteria was also included.

PCRs of 25 µL were composed of the following reagents in the following order: 1x PCR buffer (*Roche*), 260 µM of each of dATP, dCTP, dGTP, and 520 µM of dUTP (*GeneAmp, Applied Biosystems*), primermix (see table 3 for individual final concentration), 0.25 U FastStart Taq DNA Polymerase (*Roche*), 0.25U UNG (*Applied Biosystems*), 2.6 mM MgCl₂ and 5 µL template DNA.

All 18 primer sequences and their individual final PCR concentrations are shown in table 3. DNA amplifications were performed in a MJ Research (PTC-200) thermocycler, with the following program: 50 °C for 1 min, 94°C for 6 min, 35 cycles of [94°C for 50 s, 57°C for 40 s and 72°C for 50 s] and 72°C for 7 min.

Cell preparations

Bacterial templates for sensitivity studies were prepared as follows. Reference strain D2164, fr1368, D1826, D2168 and D2103 were grown to medium sized colonies (1-2 mm) on agarose plates. One of each colony was transferred to 100 µL 10 % Chelex (*Bio-rad*) and boiled for 5 min. The supernatants from the Chelex preparations were combined in the

following way. Ten μL of strain fr1368 (*ipaH* positive) and 10 μL of strain D2164 (*vtx1*, *vtx2*, *eae* and *ehxA* positive) were mixed and 10-fold serially diluted in water. Ten μL of each of these dilutions were mixed with 10 μL the D2103 (non-pathogenic strain) supernatant, from where 5 μL was used for PCR. Strains representing the other virulence genes D1826 (*eae* and *bfpA* positive) and D2168 (*sta* and *elt* positive) were prepared the same way. Bacterial templates for the specificity study, were prepared by growing reference strains on agarose plates. One of each colony was transferred to 100 μL 10% Chelex (*Bio-rad*) and boiled. Five μL of the supernatant was used directly in the PCR.

10 Detection of PCR-products:

PCR products were identified on a standard agarose gel electrophoresis system by ethidium bromide staining. Gels were made of 1.5% agarose and applied voltage was 4.5 volts/cm.

RESULTS AND DISCUSSION

15 PCR primers were designed on basis of the sequence comparisons described in section 5 (Detailed description of the invention). The amplicons representing the different virulence genes were all of unique sizes, being easily distinguished by standard agarose gel electrophoresis (table 1). Intensive specificity and sensitivity studies are often not included in papers describing diagnostic PCR analyses (17,19) It is however, very important to test both parameters thoroughly before an experimental procedure is introduced into a diagnostic laboratory.

20 Table 4. Strains used to test the sensitivity limit of the multiplex-PCR assay. Strain fr1368 and strain D2164 were mixed in equal volumes and 10-fold serially diluted relative to the non-pathogenic strain D2103, thereby testing the sensitivity limit of the *ipaH*, *vtx1*, *vtx2*, *eae* and *ehxA* genes. Strain D2168 and strain D1826 were treated likewise in order to test the sensitivity of *sta*, *elt*, *eae* and *bfpA* genes.

Strain nr.	Virulence gene(s)
fr1368	<i>ipaH</i>
D2164	<i>vtx1</i> , <i>vtx2</i> , <i>eae</i> , <i>ehxA</i>
D2168	<i>sta</i> , <i>elt</i>
D1826	<i>eae</i> , <i>bfpA</i>
D2103	non-pathogenic strains, no virulence genes

During development and optimising of the present PCR setup, a number of reference strains were tested in 10 fold serial dilutions (table 4). In order to mimic a more realistic situation 2 dilutions were made; one contains *ipaH*, *vtx1*, *vtx2*, *eae* and *ehxA* (figure 1), and another dilution contains *sta*, *elt*, *eae* and *bfpA* (figure 2). Besides the 10-fold dilutions, the pathogenic strains were also diluted in a constant volume of a non-pathogenic *E. coli* strain, mimicking a situation where very few pathogenic bacteria are present in a population of non-pathogenic bacteria. This is probably the most important way to test the sensitivity limit of PCR assays, as this allows the assay to be applied on a mixture of colonies grown from faecal sample of diarrheagenic patients. Moreover, this PCR could be applied on total DNA extracted from faeces, even if very few pathogenic bacteria are present. For both dilutions, specific amplicons were detected until a dilution of $1/10^4$ relative to D2103. If one medium sized colony is estimated to contain 10^8 bacteria and there is $1/20$ colony present in each PCR, this means that at a pathogenic strain dilution of $1/10^4$ this would correspond to a sensitivity limit at approximately 500 bacteria per PCR.

In order to test the specificity of the assay 116 reference strains obtained from The International *Escherichia* and *Klebsiella* Centre, WHO, Statens Serum Institut, Denmark, were tested by the assay.

The strains were collected over a period of 19 years (1984 to 2003), and are therefore expected to constitute a broad range of relevant clinical samples representing many different homologues of the different genes (table 5). All PCR results showed the same virulence genes as also identified by standard DNA hybridisation. Example of PCR results are shown in figure 3 and 4.

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Table 5. 116 reference strains obtained from the International *Escherichia* and *Klebsiella* Centre, WHO, Statens Serum Institut, Denmark. Before PCR, all strains were serotyped and tested for the 8 virulence genes by DNA hybridisation. When tested by the multiplex-PCR method, all strains were found positive for the exact same virulence genes as was found by hybridisation. DAEC and EAggEC were identified by probe hybridisation for other genes not included in the PCR method.

Nr.	<i>E. coli</i> Group	Year	Serotype	<i>sta</i>	<i>vtx1</i>	<i>bfpA</i>	<i>eae</i>	<i>vtx2</i>	<i>elt</i>	<i>ehxA</i>	<i>ipaH</i>	Negative for the 8 virulence genes
1	A/EEC	2003	O4, 123:H-				+					
2	A/EEC	2003	O145:H-				+					
3	A/EEC	2003	O116:H+				+			+		
4	A/EEC	2002	Oru:H8				+					
5	A/EEC	2002	O118				+					
6	A/EEC	2002	O35,135:H11				+					
7	A/EEC	2002	O51:H49				+					
8	A/EEC	2002	Oru:H33				+					
9	A/EEC	2003	O177:H25				+					
10	DAEC	1999	O21:H10							+		
11	DAEC	2000	O15:H-									+
12	EAggEC	2001	O107:H+									+
13	EAggEC	2001	O153:H2									+
14	EAggEC	2001	O92:H33									+
15	EAggEC	2001	O103:H+									+
16	EAggEC	2001	O92:H+									+
17	EAggEC	2001	O150:H28									+
18	EAggEC	2001	O24:H+									+
19	EAggEC	2001	O49:H-				+			+		+
20	EAggEC	2001	O113:H-									
21	EIEC	2000	O64:H-									
22	EIEC	2000	O64:H-								+	
23	EIEC	2001	O+-H-								+	
24	EIEC	2001	O121:H-								+	
25	EIEC	2001	O28ac:H-								+	
26	EIEC	2001	O173:H-								+	
27	EIEC	2002	O144:H-								+	
28	EIEC	2002	O+-H-								+	
29	EIEC	2002	O28ac:H-								+	
30	EPEC	2002	O103:H 2				+				+	
31	EPEC	2002	O111:H-			+	+					
32	EPEC	2002	O 26:H-				+					
33	EPEC	2002	O142:H34			+	+			+		
34	EPEC	2002	O129:H11				+					
35	EPEC	2002	O111:H38				+					
36	EPEC	2002	O111:H 9				+					
37	EPEC	2002	O114:H49				+					
38	EPEC	2002	O145:H34				+					
39	EPEC	2002	O121:H19				+					
40	EPEC	2002	O126:H6				+					
41	EPEC	2002	O 26:H -				+					
42	EPEC	2002	O 55:H 7				+			+		
43	EPEC	2002	O125ab:H 5				+					
44	EPEC	2002	O127:H-			+	+					
45	EPEC	2002	O145:H-				+			+		
46	EPEC	2003	O 26:H-				+					
47	EPEC	2003	O145:H-				+			+		

112	VTEC	1997	O145:H 4		+	+	+
113	EPEC	2001	O114:H-	+	+		
114	EPEC	2001	O 86:H 8	+	+		
115	A/EEC	2001	O132:H34	+	+		
116	EPEC	2001	O 86:H 8	+	+		

Finally, 16 non-*E. coli* strains were tested by the PCR method in order to investigate any cross reaction to other species. The strains were: *Salmonella enteritidis*, *Salmonella para A*,
5 *Salmonella typhimurium*, *Vibrio cholera*, *Aeromonas caviae*, *Shigella flexneri* 2 a, *Shigella dysenteriae* 3, *Proteus*, *Pseudomonas*, *Plesiomonas shigelloides*, *Serratia marcescens*, *Shigella sonnei*, *Klebsiella*, *Citrobacter freundii*, *Salmonella choleraesuis*, *Yersinia ent.* biotype 5, 27. Except for the 3 *Shigella* species that showed an *ipaH* band, none of the other species resulted in any signals (data not shown)

Example 2:

In this example a less complex multiplex-PCR assay was tested on 499 of clinical samples that were also tested by DNA hybridisation. This multiplex-PCR is able to identify the same
15 *E. coli* groups (VTEC, EPEC and A/EEC, ETEC or EIEC) as the method described in examples 1, but relies on fewer genes and therefore gives a less informative diagnostic answer. The PCR analysis is directed towards genes encoding the following virulence factors: heat-labile enterotoxin (LT) characteristic for ETEC, heat-stable enterotoxin (ST) characteristic for ETEC, intimin (Eae) characteristic for EPEC, A/EEC or VTEC, vero toxin 1 and 2 (VT1 and VT2) characteristic for VTEC and invasive plasmid antigen (IpaH)
20 characteristic for EIEC. The PCR analysis is performed on template DNA derived from bacterial colonies grown from faecal samples. Each PCR was analysed by agarose gel electrophoresis for the possible presence of amplicon(s) of the size that could be identified as any of the virulence markers mentioned above. In this example, the diagnostic quality of multiplex-PCR analysis is compared to membrane hybridisation, which is a traditional
25 method for diagnostics of pathogenic *E. coli* groups.

MATERIALS AND METHODS

Sample handling:

Bacterial colonies were cultured from faecal samples as follows. A small volume
30 (approximately 0.1 g) of the faeces sample was gently shaken in 2 mL sterile buffered

saltwater (80 mM NaCl, 50 mM Na₂HPO₄, 10 mM KH₂PO₄, pH 7.38). Approximately 10 µL of that suspension was streaked out onto an agar plate (*SSI Enteric Medium, Statens Serum Institut, Denmark*) and grown overnight at 37°C.

5 **Membrane hybridisation:**

As described in Example 1.

Template preparation:

10 From each clinical samples a number of morphological different colonies were picked from the plate grown cells and transferred to the same 200 µL 10 % Chelex (*Bio-Rad*) and boiled for 5 min. Five µL of supernatants were used for PCR.

PCR

15 The multiplex-PCR contained primer set for the following genes: *elt*, *sta*, *eae*, *vtx1*, *vtx2*, *ipaH* and *16S* rDNA in concentration listed in table 3. All other parameters and reagents were the same as described in example 1.

Detection of PCR-products:

20 PCR products were identified on a standard agarose gel electrophoresis system by ethidium bromide staining. Gels were made of 1.5% agarose and applied voltage was 4.5 volts/cm.

RESULTS AND DISCUSSION

25

Growing faecal samples on SSI Enteric Media allows a certain differentiation of bacterial species based on phenotypical characteristics. Different strains are however not always associated with a significant phenotype. It is therefore important to pick as many different colonies as possible from a plate, in order to increase the chance of having the possible pathogenic bacteria included. With the sensitivity limit established in example 1, a positive result can be obtained if just one of the picked colonies among 10⁴ colonies is positive.

30

Table 6. Performance of multiplex-PCR and membrane hybridisation in faeces diagnostics of *E. coli*. A total of 499 samples were analysed; both method found the same 27 samples positive and 465 samples negative. 5 samples were tested positive only by PCR, whereas 2 samples were positive only by hybridisation. All discrepancies were double tested by PCR.

Gene	Found by both methods	Only found by multiplex-PCR	Only found by DNA-hybridisation	Total
Negative	465			
Eae	19	2	1	
VT1	1	1		
VT2	1	1		
LT	3			
VT1 + eae	1			
ipaH	2	1	1	
Total	492	5	2	499

In the present example 499 clinical samples were tested by both multiplex-PCR and DNA hybridisation. The results are summarized in table 6. Both methods detected the same 465 samples as negative and 27 positive samples distributed on 6 genotypes. Five samples were found positive only by PCR, whereas 2 samples were positive only by DNA hybridisation.

All discrepancies were retested by PCR and showed the same result, decreasing the likelihood of a failed PCR. One explanation of the different result could therefore be differences in the quality of the picked colonies. As different persons might have picked different colonies and as some plates might have had very few pathogenic colonies on them, some differences are expected to be present in the DNA composition used in the 2 assays.

With the relative higher number of positives found by PCR and the obvious time saving advantages, the PCR method is concluded to be superior to the hybridisation technique.

20 . **Example 3:**

INTRODUCTION

Multiplex-PCR analysis for the diagnosis of the pathogenic *E. coli* strains; VTEC, EPEC and A/EEC, ETEC or EIEC in human faecal samples. The PCR analysis is performed on template

DNA purified directly from human stool samples. The subsequent PCR analysis is directed towards genes encoding the following virulence factors: LT characteristic for ETEC, ST characteristic for ETEC, Eae characteristic for EPEC, A/EEC or VTEC, BfpA characteristic for EPEC, EhxA characteristic for VTEC, VT1 and VT2 characteristic for VTEC and IpaH characteristic for EIEC. Completed PCRs were analysed by the Luminex® technology, where PCR products are hybridised to specific capture-probes situated on microbeads.

MATERIALS AND METHODS

DNA preparation:

DNA isolation from faecal samples was done by using the KingFisher 96™ from ThermoLabsystem, according to the manufactures instruction. Briefly, a small volume (approximately 0.1 g) of the faeces sample was gently shaken in 2 mL sterile buffered saltwater (80 mM NaCl, 50 mM Na₂HPO₄, 10 mM KH₂PO₄, pH 7.38). 200 µL of each faecal suspension was mixed with 750 µL lysis buffer and loaded in separate wells in a 96 well microtiter plate. Other plates were prepared containing either: washing buffer, 70% ethanol, distilled water and suspensions of magnetic particles. Five µL of the final DNA concentrate was used per PCR.

Sensitivity studies were performed, by adding 10-fold serially dilutions, of strains harbouring virulence gene(s), to a faecal sample tested negative for that specific gene(s).

PCR:

PCR conditions were the same as described in Example 1, except that the forward primers were 5'-biotinylated in this analysis.

Detection of PCR-products:

PCR products were detected by the Luminex® technology. Amplicons were labelled by having the forward primers 5'-biotinylated. Capture probes (listed in table 7) were synthesized with a 5'-amine Uni-Link modification, for the coupling to the carboxylated microbeads.

Each probe was coupled to a uniquely coloured population of carboxylated microbeads. This was done by mixing 1 nmol of amine-substituted probe with a suspension of 5×10^6 microbeads in 50 μ L 0.1 M 2-(N-Morpholino) ethanesulfonic acid, pH 4.5 [MES], followed by addition of 25 μ g N-(3-Dimethylaminopropyl)-N'-ethylcarbodiimide [EDC] and incubation in the dark for 30 min. The EDC addition and incubation were repeated and the microbeads were washed once with 0.02% Tween-20 and once with 0.1% SDS. Coupled microbeads were stored in TE buffer (10 mmol/L Tris-HCl, 1 mmol/L EDTA, pH 8.0) in the refrigerator in the dark until hybridization.

Five μ L of the PCRs was denatured at 95°C for 10 min and added to the hybridization solution (3 M tetramethylammonium chloride, 50 mM Tris-HCl, pH 8.0, 4 mM EDTA, pH 8.0, 0.1% sakrosyl) containing a mixture of 5000 of each probe-coupled microbead in a 50 μ L total reaction volume. Reactions were hybridized at 55°C for 10 min, pelleted by microcentrifugation and resuspended in 50 μ L hybridization solution. Hybridized amplicons were labelled with 120 ng streptavidin-R-phycoerythrin at 55°C for 5 min (62 μ L total reaction volume). Reactions were then analyzed on the Luminex® 100.

Table 7. Oligonucleotide sequence of the capture probes used in the hybridization analyses of the 9 genes. Probe design is described in "Detailed description of the invention".

Virulence factor	Encoded by gene	Oligonucleotide probe sequences (5' → 3')
Heat labile entero-toxin I (ST)	<i>elt</i>	CACAACCTTGTGGTGCATGATGAATCCAG
Heat stable entero-toxin I (LT)	<i>sta</i>	CACAGCAGTAATTGCTACTATTCATGCTTTCAGGA
Intimin (Eae)	<i>eae</i>	TACCCGTTTAGGTATTGGTGGCGAATACTGG
Verocytotoxin 1 (VT1)	<i>vtx1</i>	TCCAGAGGAAGGGCGGTTTAATAATCTACGG
Verocytotoxin 2 (VT2)	<i>vtx2</i>	TGGTTTCATCATATCTGGCGTTAATGGAGTTTCAG
Invasion plasmid antigen H (IpaH)	<i>ipaH</i>	CCAGCATCTCATACTTCTGCTCTTCTGCCTG
Enterohemolysin A (EhxA)	<i>ehxA</i>	TGCTGAGAAAACAACGGGAAGGAGAGGA
Bundle-forming pilus A (BfpA)	<i>bfpA</i>	TCAGAAGTAATGAGCGCAACGTCTGCAATT
16S rDNA	<i>16S</i>	AACGTATTCACCGTGGCATTCTGATCCAC

DISCUSSION

Primer- and probe design were performed as described in Example 1 and in "Detailed description of the invention"

- 5 KingFisher™ is an semi-automated equipment that can purify DNA from complex biological materials in about 30 minutes. The technique relies on the binding of DNA to magnetic particles that are efficiently separated and washed to produce a pure template DNA preparation. The apparatus handles 96 samples in one analytical setup by use of microtiter plates. Four microtiter plates containing different reagents need to be manually prepared
- 10 before hand. A number protocols have all ready been established for the preparation of template DNA from different starting materials. These include for example blood, different tissues and stool samples. With this flexibility in mind, the KingFisher™ is expected to perform well on most chemical complex primary sources.
- 15 Compared to growing samples, DNA purification directly from the primary source has several advantages. One, the time of analysis is reduced, as this automated DNA purification can be done in less than as hour, compared to the over night growth needed to obtain usable plate grown colonies. Two, DNA purified directly from the source should represent the different DNA populations in the same relative distribution as in the sample, including DNA
- 20 from dead bacteria. This is advantageous since *in vitro* growth will be favorable to some bacteria and make others loose their plasmids. Three, the automated procedure using this apparatus requires less hands-on-time than the growing of samples, which also reduces the tedious and repetitive manually work.
- 25 When the Luminex® technology is applied for the detection of PCR products, the analysis is based on a stringent hybridization to specific capture probes. Each specific capture probes is situated on a population of microbeads that has a distinguishable internal colour. The hybridization reaction is automatically analyzed by measuring the combination of microbead-colour and the PCR product colour-labeling, one bead at a time. Due to limitations of the
- 30 internal color-coding of the microbeads, the technology has a maximum potential of screening with 100 probes at a time. One hundred gene probes are expected to go a long way, even if other diarrheagenic bacteria like for example *Salmonella*, *Camphylobacter*, *Yersinia*

and *Klebsiella*, and genes encoding antibiotic resistance are included in the assay. In this case, the multiplex-PCRs need to be divided into a suitable number of reactions. The Luminex technology has proven very sensitive and able to discriminate between single mutations in SNP analysis (29), and is therefore expected to enclose the analytical capacity necessary for the present diagnostic assay.

Compared to gel electrophoresis the Luminex technology has a number of advantages: 1) the sequence specific hybridization adds more specificity to the diagnostic assay, compared to the un-precise size determination of gel electrophoresis, 2) the Luminex procedure is faster, semi-automated and therefore requires less hands-on-time 3) expansion of gene number would be limited in gel electrophoresis as each gene has to have a distinguishable size, whereas Luminex will screen for up to 100 genes in the same reaction.

Example 4:

INTRODUCTION

Multiplex-PCR analysis for the diagnosis of the diarrheagenic *E. coli* strains; VTEC, EPEC and A/EEC, ETEC or EIEC in human faecal samples. The PCR analysis is performed on template DNA purified directly from human stool samples, by use of BUGS'n BEADS™ from Genepoint A.S. (Oslo, Norway). The subsequent PCR analysis is directed towards genes encoding the following virulence factors: LT characteristic for ETEC, ST characteristic for ETEC, Eae characteristic for EPEC, A/EEC or VTEC, BfpA characteristic for EPEC, EhxA characteristic for VTEC, VT1 and VT2 characteristic for VTEC and IpaH characteristic for EIEC. Completed PCRs were analyzed by the hybridization to specific capture probes that were immobilized on plastic surfaces (PCR-ELISA) and microarrays.

MATERIALS AND METHODS

DNA preparation:

A small volume (approximately 0.1 g) of the faeces sample was gently shaken in 2 mL sterile buffered saltwater (40 mM NaCl, 25 mM Na₂HPO₄, 5 mM KH₂PO₄, pH 7.38). This suspension was left standing for 2 min to obtain a supernatant free from most undissolved materials. The DNA was isolated by use of the BUGS'n BEADS™ from Genepoint A.S. (Oslo, Norway). 500 µL supernatant for the sample was mixed with 400 µL Binding Buffer

and 20 μ L Bacterial Binding Beads in a 1.5 mL microcentrifuge tube. The bead/bacterial complex was held on the inside of the tube, by using an external magnet, while the supernatant was poured off. The bead/bacterial complex was dissolved in 50 μ L Lysis Buffer and incubated at 80°C. After 5 min 150 μ L 96% cold ethanol was added and the incubation was continued for another 5 min. The bead/bacterial complex was retained by magnetic force while the supernatant was discharged. The bead/bacterial complex was washed twice with 70% ethanol. The bead/bacterial complex was finally resuspended in 30 μ L H₂O, from where 5 μ L was used for PCR.

Probes for both ELISA hybridization and microarrays are the same. Probe- and primer design was based on the criteria describes in "Detailed description of the invention".

ELISA hybridisation

The attachment of the capture probes to the solid surface and the hybridizations were done with NucleoLink™ Surface (PCR-ELISA) from Nalge Nunc International according to the manufactures instructions. Oligonucleotide probes for each of the 9 genes are listed in table 7. Each probe was synthesized with 12 T's added to the 5'-end, and the final T in the 5'-end being phosphorylated. The 12 T's served as a linker that increases the hybridization efficiency by lifting the probe away from the solid surface. The phosphorylation was necessary for the covalent binding of the probe to the plastic surface. Approximately 100 nM of each of the 9 capture probes were used in the covalent binding to the surface of separate wells in a 96-well microtiter plate. The PCR was performed as described in Example 1 except that primers were biotinylated at the 5'-end and the total reaction volume was 100 μ L. The hybridization was done by adding 10 μ L of the PCR to each of the 9 wells containing the different capture probes. After a standard hybridization procedure, including NaOH denaturation, hybridization at 52°C for 2 h and SSC incubations and washing, the reaction was detected by use of Alkaline phosphatase conjugated streptavidin (DAKO, Denmark).

Microarray detection

Primers used in the PCR were Cy3-labelled in the 5'-end, and the total reaction volume was 100 μ L. All other PCR parameters were the same as described in Example 1. Completed

PCRs were purified with the QIAquick PCR Purification kit from QIAGEN, dried and resuspended in 4 μL H_2O , 7.5 μL 20 x SSC, 2.5 μL 1% SDS, 1 μL salmon sperm DNA (10 mg/mL) and 15 μL formamide. This suspension was denatured at 94°C for 5 min prior to the hybridisation.

5

Capture probes used in this experiment are listed in table 7. Approximately 3 nmol of each probes was dissolved in 20 μL 6 N Na_2SCN . One nL of the probes were arrayed onto coated glass slides (Amersham Pharmacia Biotech) and baked at 80°C for 2 h. Prior to pre-hybridization the slides were washed in isopropanol and boiling water and dried in ultra clean N_2 . The pre-hybridization consisted of 20 min incubation at 60°C in 3.5 x SSC, 0.2% SDS and 1% BSA, rinsing in water and isopropanol and drying in ultra clean N_2 gas. Thirty μL of the denatured sample was added to the pre-hybridized slide and covered with a coverslip and stored in a moisturized hybridization chamber for 14 h at 35°C. The coverslip was removed and the slide was washed 5 times in SSC and 0.1% SDS, SSC sequentially being 2x, 1x, 0.1x, 0.1x and 0.1x. The slides were finally dried in ultra clean N_2 . The hybridization signal was analysed by a confocal laser microscope (Molecular Dynamics, California).

20

DISCUSSION

Bugs n' BeadsTM is a simple and relative inexpensive procedure for the preparation of bacterial DNA from growth media and biological samples. It is designed, first to separate bacteria by their adherence to magnetic beads, and then to prepare DNA from the isolated bacteria. The bacterial-bead extraction is a powerful preparation technique, as it separates bacteria from all non-bacterial materials in one single step. Also, if the cell count of the sample is low, this concentrating procedure will help to obtain a detectable signal. The technique is therefore expected to perform well on chemical complex samples as for example faeces or foods. The relative simple procedure of Bugs n' BeadsTM is well suited for automatisisation, which would be valuable for a routine diagnostic laboratory.

30

ELISA hybridization is a relative established method, which therefore has the advantages of being less prone to technical problems. Each gene must be hybridized in separate tubes/wells,

which makes the procedure more tedious than Luminex, where all hybridizations can take place in the same reaction tube.

5 Microarrays has the advantage of being extremely well suited for screening of many genes in the same sample. This has been shown usable in many studies dealing with for examples the expression of thousands of genes. In the case of analyzing many samples for the existence of a few genes, microarrays are relative expensive. The technique is therefore expected to be most relevant when a more complete assay is undertaken, dealing with many different strains, antibiotic resistance and subtyping. With the proper design, such a “universal pathogen
10 microarray” would find use in many places and eventually on the commercial marked at a lower price.

Compared to gel electrophoresis, both ELISA and microarrays have the following advantages 1) the sequence specific hybridization adds more specificity to a diagnostic assay
15 compared to the un-precise size determination of gel electrophoresis, 2) the Luminex procedure is faster, semi-automated and therefore requires less hands-on-time 3) expansion of gene number would be limited in gel electrophoresis as each gene has to have a distinguishable size, whereas Luminex will screen for up to 100 genes in the same reaction.

20 Example 5

INTRODUCTION

Multiplex-PCR analysis for the diagnosis of the pathogenic *E. coli* strains; VTEC, EPEC and
25 A/EEC, ETEC or EIEC in human faecal samples. The real-time PCR analysis was performed on template DNA purified directly from human stool samples by use of FTA Technology from Whatman. Genes encoding the following virulence factors were targeted: LT characteristic for ETEC, ST characteristic for ETEC, Eae characteristic for EPEC, A/EEC or VTEC, BfpA characteristic for EPEC, EhxA characteristic for VTEC, VT1 and VT2
30 characteristic for VTEC and IpaH characteristic for EIEC.

MATERIALS AND METHODS

DNA preparation:

The DNA extraction was performed by FTA® Technology from Whatman. A small volume (approximately 0.1 g) of the faeces sample was gently shaken in 2 mL sterile buffered saltwater (40 mM NaCl, 25 mM Na₂HPO₄, 5 mM KH₂PO₄, pH 7.38). This suspension was left standing for 2 min to obtain a supernatant free from most undissolved materials. Ten µL of that supernatant was transferred to the FTA Card and allowed to dry. A small circle of two mm in diameter of the card containing the sample was punched out and put into a 0.5 mL PCR tube. The punch was first washed three times in 200 µL FTA Purification Reagent and then washed twice in TE. After the final wash the punch was allowed to air dry and used directly as template in the PCR amplification.

Real-time PCR:

Due to fluorescence overlap and technical equipment limitations, real-time multiplex PCR is presently limited to analyze four genes in the same reaction. Therefore, the total analysis of 9 genes needed to be performed in three separate multiplex reactions. The combinations of genes, primers, probes, fluorophores and quenchers in each of the three reactions are listed in table 8. The primers used in this analysis are the same as in the previous examples (see table 3 for primer sequences and table 7 for probe sequences). In each reaction the different probes were coupled to fluorophores with distinguishable emission spectra, and a compatible quenchers. The analysis was performed on a Mx4000™ Multiplex Quantitative PCR System from Stratagene.

Table 8. Real time PCR parameters. See table 2 and 5 for primer and probe sequences.

Reaction number	Genes targeted	Forward / reverse primer	Forward / reverse primer concentration (μM)	Probe labelling: 5'-fluorophore / 3'-quencher	Probe concentration (μM)
1	<i>sta</i>	ST-F / ST-R	0.3 / 0.3	5-FAM / TAMRA	0.2
	<i>vtx1</i>	VT1-F / VT1-R	0.25 / 0.25	5-ROX / BHQ-2	0.15
	<i>ehxA</i>	Ehx-F / Ehx-R	0.15 / 0.15	Cy5 / BHQ-3	0.2
	<i>16S</i>	16S-F / 16S-R	0.25 / 0.25	HEX / BHQ-1	0.1
2	<i>elt</i>	LT-F / LT-R	0.4 / 0.4	5-FAM / TAMRA	0.15
	<i>eae</i>	Eae-F / Eae-R	0.2 / 0.2	5-ROX / BHQ-2	0.2
	<i>ipaH</i>	IpaH-F / IpaH-R	0.1 / 0.1	Cy5 / BHQ-3	0.2
	<i>16S</i>	16S-F / 16-R	0.25 / 0.25	HEX / BHQ-1	0.1
3	<i>bfpA</i>	Bfp-F / Bfp-R	0.2 / 0.2	5-FAM / TAMRA	0.2
	<i>vtx2</i>	VT2-F / VT2-R	0.25 / 0.25	5-ROX / BHQ-2	0.15
	<i>16S</i>	16S-F / 16-R	0.25 / 0.25	Cy5 / BHQ-3	0.15

BHQ: balck hole quencher

Each PCR parameter was tested at different values in order to find optimum condition that resulted in highest sensitivity and specificity of every gene in the assay. The total reaction volume was 25 μL and contained the following reagents: 1x PCR buffer (Applied Biosystems), 260 μM of each of dATP, dCTP, dGTP and 520μM og dUTP (Applied Biosystems), primer- and probe- combinations and concentrations as listed in table xx. 0.5U AmpliTaq Gold® DNA Polymerase (Applied Biosystems), 0.25U UNG (Applied Biosystems), 3.2 mM MgCl₂ and 5 μL template. The temperature parameters were as follows: 2 min at 50°C, 10 min at 94°C and 40 cycles of 15 sec at 95°C and 1 min at 58°C. Data analysis was done by the software included in the Mx4000™ Multiplex Quantitative PCR System.

DISCUSSION

FTA technology from Whatman is based on cell lysis and DNA entrapment in the fibrous matrix of a specialized membrane. In the subsequent washing steps the cell debris and PCR inhibitors are removed. Finally, a small piece of the membrane is added directly to the PCR from where the DNA still bound to the matrix serves as a template for the amplification reaction. The FTA technology is expected to perform well in the DNA purification from

many different starting materials, including stool samples. This is based on the fact, that the DNA is bound relative stable early in the purification procedure to the membrane, and that this allows an intense washing of the DNA-membrane complex, resulting in an efficient removal of PCR inhibitors.

5

Primers and probes are designed as described in “detailed description of the project” and in Example 1. The real-time-multiplex analysis is restricted to four simultaneous reactions per reaction tube because of the overlap in emission spectra. However, as the 9-gene multiplex PCR analysis has been optimized to work well under non real-time PCR conditions, it is
10 believed that when the technology is developed to contain more fluorophores in the same reaction, it should be possible to use the present PCR setup in a “9-gene-multiplex- real-time-PCR” setup. Until then, the 9-gene analysis is divided into 3 reaction-tubes reaction which is still an reasonable analytical strategy.

15 **Example 6**

INTRODUCTION

Multiplex-PCR analysis for the diagnosis of the pathogenic *E. coli* strains; VTEC, EPEC, ETEC or EIEC in human faecal samples. The PCR analysis is performed on template DNA purified directly from human stool samples by using the QIAamp® DNA Stool Mini Kit (QIAGEN). The subsequent PCR analysis is directed towards genes encoding the following virulence factors: LT characteristic for ETEC, ST characteristic for ETEC, Eae characteristic for EPEC, A/EEC or VTEC, BfpA characteristic for EPEC, EhxA characteristic for VTEC, VT1 and VT2 characteristic for VTEC and IpaH characteristic for EIEC. Completed PCRs
20 were analysed by capillary electrophoresis.

25

Materials and methods:

DNA preparation:

30 The DNA was purified by QIAamp® DNA Stool Mini Kit from QIAGEN according to the provided instructions. Briefly, 0.1-0.2 g of faeces was measured into a 2 mL microcentrifuge tube and mixed with 1.4 mL Buffer ASL. This suspension was incubated at 70°C for 5 min, vortexed for 15 s and centrifuged for 1 min. The supernatant was then treated with InhibitEX that binds PCR inhibitors and precipitates them during a 3 min of centrifugation step at

13.000 g. 600 μ L of the supernatant was then treated with 25 μ L Proteinase K, mixed with Buffer AL, vortexed for 15 s and incubated at 70°C for 10 min. The lysate was mixed equal volumes of 96% ethanol spun in QIAamp spin columns. The bound DNA was washed by first running 500 μ L Buffer AW1 and secondly, by running 500 μ L Buffer AW2 through the column. The DNA was eluted by incubating the column with 200 μ L Buffer AE for 1 min and spinning the column at 13.000g for 1 min.

PCR:

PCR conditions were the same as described in Example 1, except that the forward primers were synthesized containing 5-FAM on the 5'-prime end.

Detection of PCR-products:

The PCR products were analyzed by capillary electrophoresis on an ABI PRISM 310 Genetic Analyzer (Perkin-Elmer) according to the manufactures instructions. The capillary was 47 cm by 50 μ m, the temperature was constant at 60°C, and the electrical field was 15 kV. One microliters of the completed PCRs and 1 μ L 2500-TAMRA labelled size standard were denatured by incubating in 12 μ L formamide at 95°C for 3 min, and cooled on ice before loading on the analyser. The resulting peaks (5-FAM labelled) were identified by comparison to the size standard.

DISCUSSION

QIAamp® DNA Stool Mini Kit from QIAGEN is developed to purify DNA directly from stool samples, either with increased ratio of non-human to human DNA or vice versa depending on the analytical need. The kit relies on bacterial lysis, specific absorption and precipitation of PCR inhibitors and DNA binding to washable centrifuge columns. If the centrifugation step is exchanged with a vacuum manifold, the whole procedure can be automated making it ideal for a routine laboratory.

Capillary electrophoresis is seen as a fast and more precise alternative to gel electrophoresis. The time saving depends on sample capacity of the apparatus and the number of samples to

be analyzed. If hundreds of samples are to be analyzed the gel electrophoresis might be faster if big gels with high sample number capacity are used. When it comes to size determination, capillary electrophoresis is clearly superior to gel electrophoresis, offering an overall more specific assay.

5

Example 7

INTRODUCTION

10 This example deals with the subtyping of the *E. coli* virulence genes: *vtx1*, *vtx2* and *eae*. The severity of the clinical manifestations of *vtx2* and/or *eae* infections, are believed to be at least partly determined by the specific subtype (5). The specific subtype should be taken into consideration when the proper treatment is determined. Subtyping of these important virulence markers, is also valuable for epidemiological surveillance. Based on sequence data,
15 the *eae* genes can be divided into at least 7 subtypes, whereas more than 6 subtypes of *vtx2* have all ready been identified. Many subtyping studies are based on the detection of fragment patterns of PCR products being digested by restriction enzymes (1,2,25). Unfortunately, such methods are based on relative few sequence characteristics, and restricted to include sequence spots to where suitable restriction enzymes can be found. Also, these methods can
20 be misleading, if a new mutation occurs at a site where the enzyme was designed to cut. Direct sequencing of PCR product by standard sequencing reaction, will easily yield 500 readable base pairs, which if chosen right can be very discriminating between different subtypes. As the cost of sequencing reactions, have been much reduced over the last couple of years, this method is now feasible for routine diagnostic analyses.

25

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Claims

1. A screening method for simultaneous detection of diarrheagenic *Shigella spp.* and *E. coli* (DEC) including A/EEC & EPEC, ETEC, VTEC, EIEC and strains with the *ehxA* gene.
2. A screening method according to claim 1, which detects the presence of the genes *ehxA*, *eae*, *vtx1*, *vtx2*, *ipaH*, *sta*, *elt* and *bfpA*.
3. A screening method according to claim 1-2 for detecting *Shigella spp.* by detecting the presence of the *ipaH* gene.
4. A screening method according to claim 1-3 performed with multiplex PCR.
5. A screening method according to claim 4, using the primers from table 3.
6. A screening method according to claim 4-5, which additionally incorporates a positive control.
7. A screening method according to claim 6, where the positive control is 16S rDNA.
8. A screening method according to claim 4-7, which uses the UNG system.
9. A screening method according to any of the preceeding claims where the genes are detected by size identification, e.g. by agarose gel electrophoresis or capillary electrophoresis.
10. A screening method according to any of the preceeding claims where the genes are detected with a hybridisation probe.
11. A screening method according to any of the preceeding claims where the material to be analysed can be any material from where bacteria can be extracted, e.g stool samples, consumables etc.
12. An *in vitro* diagnostic method for determining the risk of being infected with a pathogenic organism giving rise to haemolytic uremic syndrome (HUS) or hemorrhagic colitis, said method comprising detecting the *ehxA* gene in the DEC.
13. A method for simultaneously detection of diarrheagenic *E. coli* (DEC) groups A/EEC & EPEC, ETEC, VTEC, and EIEC by testing for the presence of the genes: *ipaH*, *eae*, *ehxA* and *sta*, parts of these genes or products of these genes or parts thereof, such as RNA or polypeptides.
14. A method according to claim 13, which further comprises (simultaneously) testing for the presence of one or more (such as more than 2, 3 or 4) of the genes selected from the group comprising: *vtx1*, *vtx2*, *elt*, and *bfpA*, parts of these genes or products of these genes or parts thereof, such as RNA or polypeptides.

15. A method according to claim 14, detecting the genes selected from the group comprising: *ipaH*, *eae*, *ehxA*, *sta*, *vtx1*, *vtx2*, *elt*, and *bfpA*, parts of these genes or products of these genes or parts thereof, such as RNA or polypeptides.
- 5 16. A method according to claim 14, detecting the genes selected from the group comprising: *ipaH*, *eae*, *sta*, *vtx1*, *vtx2*, and *elt*, parts of these genes or products of these genes or parts thereof, such as RNA or polypeptides.
- 10 17. A method of any of claims 13-16, in which the testing is carried out on a sample, such as a sample from a human or an animal (ie. stool), a sample from a consumable products (ie. food and beverages), a bacteria culture or a sample from sewage.
- 15 18. A method of any of claims 13-17, in which the testing is carried out using a nucleotide sequence amplification technique, such as PCR, multiplex PCR, real-time PCR.
- 20 19. A method according to any of the preceding claims, in which at least one primer used is selected from the group consisting of:
a) the primers of table 3,
b) sequences having a sequence identity of at least 80% (such as a least 85%, at least 90%, or at least 95%) with the primer sequences of a)
c) parts of the sequences in a) or b), having a length of more than 10, preferred more than 13 nucleotides (eg. consisting of 14, 15, 16, 17, 18, 19, 20, 21 or 22 consecutive nucleotides),
25 d) sequences comprising a sequence in a), b) or c), said sequence having a length of at least most 100 nucleotides, such as at most 90, 80, 70, 60, 50, 40, or at most 30 nucleotides.
- 30 20. A method according to any of the preceding claims, which additionally incorporates a positive control.
21. A method according to any of the preceding claims, where the positive control is 16S rDNA.
- 35 22. A method according to any of the preceding claims, which uses the UNG system.
23. A method according to any of the preceding claims, wherein the genes are detected by size identification, e.g. by agarose gel electrophoresis or capillary electrophoresis.
- 40 24. A method according to any of the preceding claims, wherein at least one product of the nucleotide sequence amplification reaction is detected with a hybridisation probe.
- 45 25. A nucleotide sequence selected from the group consisting of:
a) the primer sequences of table 3,
b) sequences having a sequence identity of at least 80% (such as a least 85%, at least 90%, or at least 95%) with the primer sequences of a)
c) parts of the sequences in a) or b), having a length of more than 10, preferred more than 16 nucleotides, such as more than 17, 18, 19 or 20 nucleotides (eg. consisting of

14, 15, 16, 17, 18, 19, 20, 21 or 22 consecutive nucleotides of the sequences in a) or b)),

d) sequences comprising a sequence in a), b) or c), said sequence having a length of at least most 100 nucleotides, such as at most 90, 80, 70, 60, 50, 40, or at most 30 nucleotides.

26. A nucleotide sequence, which is selected from the primer sequences of table 3.

27. A nucleotide sequence selected from the group consisting of:

a) the probe sequences of table 7,

b) sequences having a sequence identity of at least 80% (such as at least 85%, at least 90%, or at least 95%) with the primer sequences of a)

c) parts of the sequences in a) or b), having a length of more than 10, preferred more than 16 nucleotides, such as more than 17, 18, 19 or 20 nucleotides (eg. consisting of 14, 15, 16, 17, 18, 19, 20, 21 or 22 consecutive nucleotides of the sequences in a) or b)),

d) sequences comprising a sequence in a), b) or c), said sequence having a length of at least most 100 nucleotides, such as at most 90, 80, 70, 60, 50, 40, or at most 30 nucleotides.

28. A nucleotide sequence which is selected from the probe sequences of table 7.

29. A kit which comprises, in a single or in separate containers, nucleotide sequences which are able to prime amplification in a nucleotide sequence amplification reaction, such as PCR, of the genes: *ipaH*, *eae*, *ehxA*, and *st*, or parts of these genes or the complementary strands to the genes or parts thereof.

30. A kit which comprises, in a single or in separate containers, nucleotide sequences which are able to hybridise (preferably under stringent conditions) with the genes: *ipaH*, *eae*, *ehxA*, and *sta*, parts of these genes or the complementary strands to the genes or parts thereof.

31. A kit according to any of the preceding claims, which comprises at least one (such as 2, 3, 4, 5, 6, 7, 8, 9, 10 or more) nucleotide sequence of claim 25-26.

32. A kit according to any of the preceding claims, which comprises a means for detection, such as a probe with a nucleotide sequence according to claim 27-28.

33. A kit according to any of the preceding claims, which comprises a means for amplification of a nucleotide sequence, such as a polymerase or nucleotides.

34. A kit according to any of the preceding claims, which comprises a means for a control, such as primers for 16S rDNA.

35. A kit according to any of the preceding claims, which comprises a means for detecting by size identification, ie. an agarose gel or a capillary tube optionally filled with buffer.

- 5
36. A kit according to any of the preceding claims, which comprises nucleotide sequences which are able to prime amplification of at least one gene selected from the group consisting of : *vtx1*, *vtx2*, *elt*, and *bfpA*, or parts of these genes or the complementary strands to the genes or parts thereof.
- 10
37. A kit according to claim 36, in which the nucleotide sequences for priming are selected from the group consisting of the priming sequences in table 3.
38. A kit according to claim 36, in which the nucleotide sequences for probing are selected from the group consisting of the probe sequences in table 7.

Abstract

The present invention contains a method for the identification of the diarrheagenic *E. coli* groups: ETEC (enterotoxigenic *E. coli*), A/EEC (attaching and effacing *E. coli*), EPEC (enteropathogenic *E. coli*), VTEC (verocytotoxin producing *E. coli*) and EIEC (enteroinvasive *E. coli*), and *Shigella* spp. The bacterial identification is made possible by the specific detection of the following virulence genes: *stx* and *elt* encoding heat stable enterotoxin (ST) and heat labile enterotoxin (LT) characteristic of ETEC, *eae* encoding intimin, characteristic of A/EEC, EPEC or VTEC, *bfpA* encoding bundle forming pilus (BfpA), characteristic of EPEC, *vtx1* and *vtx2* encoding verocytotoxin 1 and 2 (VT1 and 2) characteristic of VTEC, *ipaH* encoding invasive plasmid antigen H (IpaH) characteristic of EIEC and *Shigella* spp., and *ehxA* encoding enterohemolysin (EhxA) characteristic of some EPEC and VTEC strains. The method allows the simultaneous detection of any combination of the 8 virulence genes by one single multiplex-PCR. The method is thoroughly validated with respect to sensitivity and specificity, and showed high performance compared to other publication. The method includes an internal positive PCR control and the carry-over prevention system, UNG, which makes it ideal for routine diagnostic analyses. The method can be combined with a number of other technologies leading to even higher sensitivity and reduced time of analysis – both important parameters when diarrheagenic patient or contaminated foods are analysed.

Figure 1

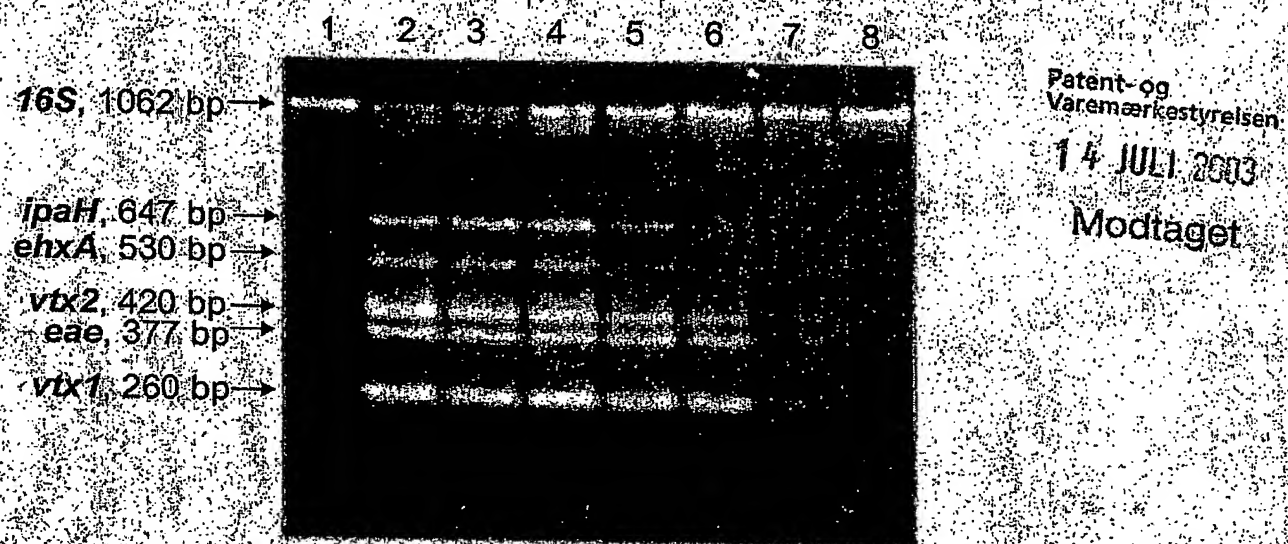


Figure 2

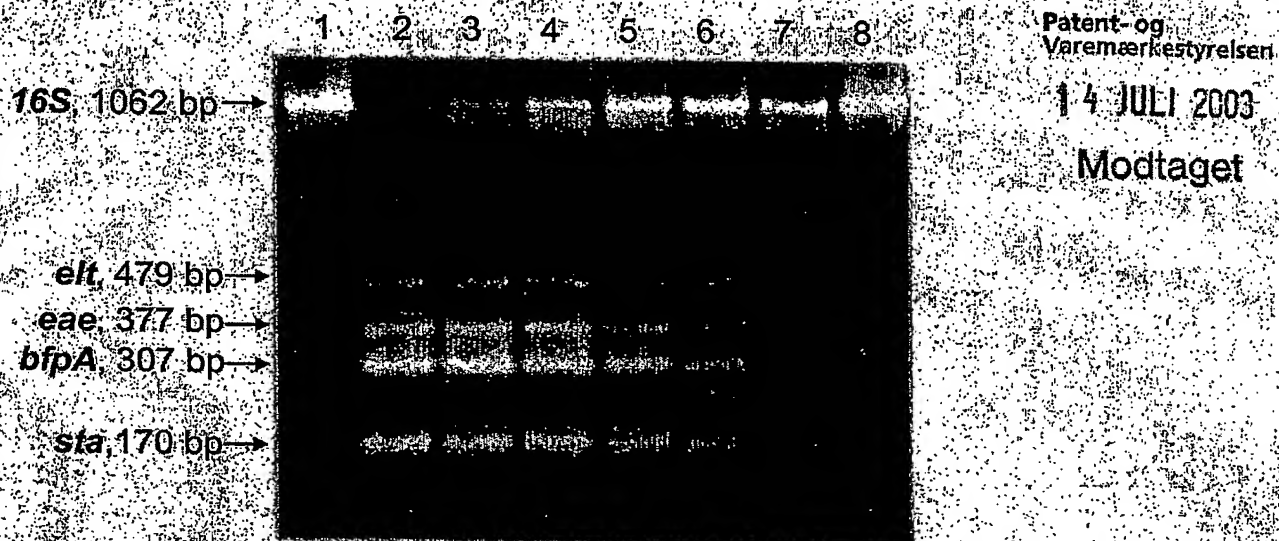
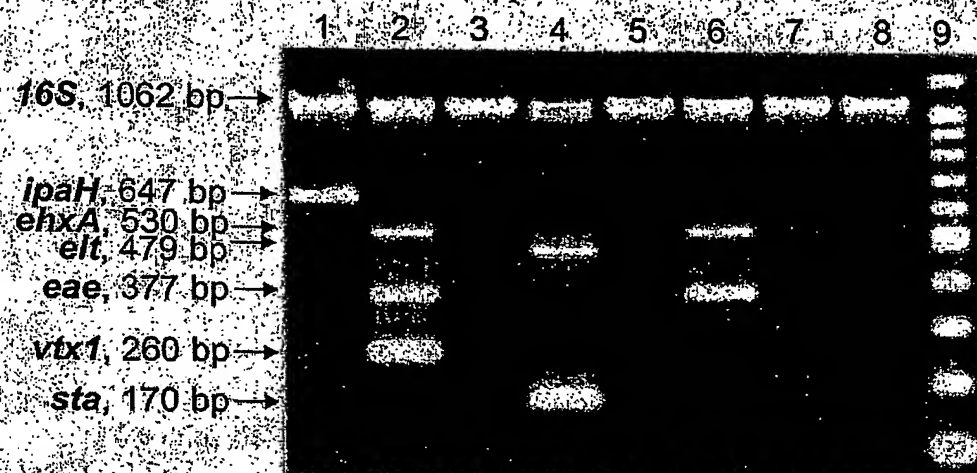


Figure 3

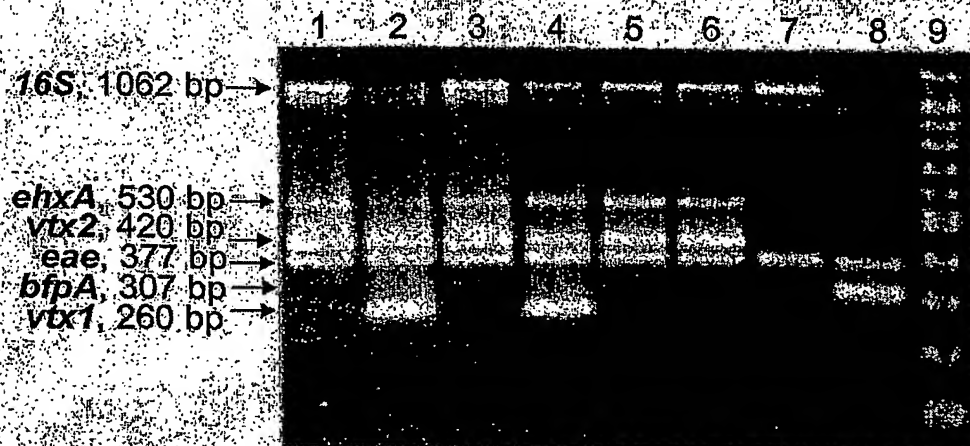


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Modtaget

Figure 4



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